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AMENDMENTS TO THE CLAIMS

This listing of claims will replace all prior versions and listings of claims in the

application:

LISTING OF CLAIMS:

1. (currently amended): A sustained release formulation for oral administration of

an HMG-CoA reductase inhibitor comprising:

a spray-dried solid dispersant in the form of particles having a particle size ranging from

5 to 200µm, wherein the solid dispersant contains containing the HMG-CoA reductase

inhibitor, a solubilizing agent, and a stabilizing agent;

a mixture of sodium alginate and xanthan gum as a sustained release composite carrier;

and

a mixture of propylene glycol ester alginate and hydroxypropyl methyl cellulose as a gel

hydration accelerator.

2. (currently amended): The sustained release formulation of claim 1, wherein the

amount of the solubilizing agent is 0.05 to 20 weight part; the amount of the stabilizing agent is

0.01 to 0.1 weight part; the amount of the sustained release composite carrier is 3 to 30 weight

part; and the amount of the gel hydration accelerator is 0.1 to 5 weight part based on 1 weight

part of the HMG-CoA reductase inhibitor.

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(original): The sustained release formulation of claim 1, wherein the HMG-CoA

reductase inhibitor is selected from the group consisting of mevastatin, lovastatin, pravastatin,

lactone of pravastatin, velostatin, simvastatin, rivastatin, fluvastatin, atorvastatin, cerivastatin

and a pharmaceutically acceptable salt thereof.

4. (original): The sustained release formulation of claim 3, wherein the HMG-CoA

reductase inhibitor is simvastatin or a pharmaceutically acceptable salt thereof.

5. (original): The sustained release formulation of claim 1, wherein the

solubilizing agent is selected from the group consisting of d- $\!\alpha\!$  -tocopheryl polyethylene glycol

1000 succinate, polyoxyethylene stearic acid ester, polyethylene glycol and polyoxypropylene-

polyoxypropylene block copolymer.

6. (original): The sustained release formulation of claim 1, wherein the stabilizing

agent is selected from the group consisting of butylated hydroxy toluene, butylated hydroxy

anisol, erythorbic acid and ascorbic acid.

(original): The sustained release formulation of claim 1, wherein the solid

dispersant further includes a pharmaceutically acceptable solubilizing carrier.

(canceled):

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9. (previously presented): The sustained release formulation of claim 1, wherein

the sustained release composite carrier includes 0.1 to 10 weight part of the xanthan gum based

on 1 weight part of the sodium alginate.

10. (previously presented): The sustained release formulation of claim 1, wherein

the sustained release composite carrier further includes locust bean gum.

11. (original): The sustained release formulation of claim 10, wherein the sustained

release composite carrier includes 0.1 to 5 weight part of the locust bean gum based on 1 weight

part of the sodium alginate.

12. (canceled).

13. (previously presented): The sustained release formulation of claim 1, wherein

the gel hydration accelerator includes 0.05 to 20 weight part of the propylene glycol ester

alginate based on 1 weight part of the hydroxypropyl methyl cellulose.

14. (original): The sustained release formulation of claim 13, wherein the

hydroxypropyl methyl cellulose has a viscosity ranging from 4,000 to 100,000 cps.

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15. (original): The sustained release formulation of claim 1, further comprising a

pharmaceutically acceptable additive selected from the group consisting of a binder, a

lubricating agent, a sweetening agent and an excipient.

16. (withdrawn): A method for preparing the sustained release formulation of claim

1, comprising the steps of:

(1) mixing the HMG-CoA reductase inhibitor, the solubilizing agent, and the

stabilizing agent in a solvent to obtain the solid dispersant;

(2) homogeneously mixing the sustained release composite carrier and the gel

hydration accelerator with the solid dispersant to form a first mixture;

(3) adding a pharmaceutically acceptable additive to the first mixture to form a

second mixture; and

(4) dry-mixing and formulating the second mixture into a solid formulation.

17. (withdrawn): The method of claim 16, wherein the solid dispersant is prepared

by a method selected from the group consisting of a spray-drying method, a solvent evaporation

method, a pulverizing wet method, a melting method and a freeze-drying method.

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18. (new): A sustained release formulation for oral administration of an HMG-CoA

reductase inhibitor comprising:

a spray-dried solid dispersant in the form of particles having a particle size ranging from

5 to 200µm, wherein the solid dispersant consists of the HMG-CoA reductase inhibitor, a

solubilizing agent, and a stabilizing agent;

a mixture of sodium alginate and xanthan gum as a sustained release composite carrier;

and

a mixture of propylene glycol ester alginate and hydroxypropyl methyl cellulose as a gel

hydration accelerator.